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Nylon 11/Silica Nanocomposite Coatings Applied by the HVOF Process. II. Mechanical and Barrier Properties

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> ABSTRACT: Nylon 11 coatings filled with nominal 0-15 vol % of nanosized silica or carbon black were produced using the high velocity oxy-fuel combustion spray process. The scratch and sliding wear resistance, mechanical, and barrier properties of nanocomposite coatings were measured. The effect of powder initial size, filler content, filler chemistry, coating microstructure, and morphology were evaluated. Improvements of up to 35% in scratch and 67% in wear resistance were obtained for coatings with nominal 15 vol % contents of hydrophobic silica or carbon black, respectively, relative to unfilled coatings. This increase appeared to be primarily attributable to filler addition and increased matrix crystallinity. Particle surface chemistry, distribution, and dispersion also contributed to the differences in coating scratch and wear performance. Reinforcement of the polymer matrix resulted in increases of up to 205% in the glass storage modulus of nanocomposite coatings. This increase was shown to be a function of both the surface chemistry and amount of reinforcement. The storage modulus of nanocomposite coatings at temperatures above the glass transition temperature was higher than that of unfilled coatings by up to 195%, depending primarily on the particle size of the starting polymer powder. Results also showed that the water vapor transmission rate through nanoreinforced coatings decreased by up to 50% compared with pure polymer coatings. The aqueous permeability of coatings produced from smaller particle size polymers (D-30) was lower than the permeability of coatings produced from larger particles because of the lower porosities and higher densities achieved in D-30 coatings. © 2000 John Wiley & Sons, Inc. J Appl Polym Sci 78: 2272-2289, 2000

Key words: polymer nanocomposite; HVOF; thermal spray; nylon 11 coatings; wear testing

INTRODUCTION

Reinforced polymer coatings used for corrosion protection are critical in applications in which increased mechanical properties and high scratch and wear resistance are required. The properties are especially enhanced by the use of nanoscalar ceramic reinforcements. Development and application of nanoreinforced polymer coatings is challenging for environmental and safety reasons be cause the use of volatile organic compounds and high processing temperatures must be avoided. In this work, the novel thermal spray processing an nanocomposite polymers was investigated. The results of scratch and sliding wear resistance aqueous permeability, and dynamic mechanical analysis of thermal sprayed nanocomposite coatings are presented. The improvements in proper

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ties were related to the filler content, filler surface chemistry, filler/matrix interactions, coating microstructure, and polymer crystallinity.

Relevant theoretical models for the prediction of nanocomposite properties including modulus, damping, and permeability are introduced. The calculated predictions are compared with experimental results and the results used to elucidate possible reinforcing mechanisms in the sprayed nanocomposite coatings.

BACKGROUND

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The inclusion of nanoparticulates in thermoplastics has been shown to result in substantial changes in polymer properties, even compared with the effects of adding micron-sized reinforcements. 1-9 Sumita et al. 1 demonstrated that 7-nm silica reinforcements in semicrystalline nylon 6 increased the yield stress by 30% and Young's modulus by 170% compared with pure polymers, phereas micron-sized reinforcements decreased he yield stress and only marginally increased the modulus. They observed similar effects for nanonized fillers in polypropylene where the dynamic storage modulus increased by 100% with the addition of 20 wt % of 7-nm silica particles whereas he addition of the same amount of 200-μm glass particles caused only a marginal increase in the dynamic storage modulus.2 The improvement in properties of layered nanocomposites was even gere pronounced because of the high aspect ratio the fillers. The dynamic storage modulus of a mnocomposite containing as low as 4 vol % of mica-type silicate in an epoxy matrix increased by in the region below the glass transition temperature (T_g) and by 450% in the rubbery region.³ ermeability of water in poly(ϵ -caprolactone) also decreased by an order of magnitude with the addition of 4.8 vol % of mica-type silicate. A 60% rease in permeability in polyimide composites ontaining as little as 2% mica was reported, mereas the thermal expansion coefficient was duced by 25%.5

The achievement of a homogeneous distribution of nanosized fillers represents an important edinical challenge in their processing because reasing filler particle size causes a significant rease in melt viscosity. Large amounts of solution of the size cause and the size causes are interest. The methods currently to produce layered nanocomposites are in-

tercalation (inserting) of the polymer between the host silicate layers or delamination of the silicate layers and dispersing them within a polymer matrix for which the intercalating monomer or polymer was also dissolved in the solvent. Other methods include *in situ* polymerization, and solgel processing for polyamide-silica nanocomposites. Recently, some progress in melt processing of mica-type nanocomposites has been reported. 9

An excellent solution to the processing limitations of polymer/ceramic nanocomposites, such as the use of solvents, is thermal spraying. During a thermal spray process, polymer particles are heated in a thermal jet (created either by a plasma or via combustion), and then accelerated toward the substrate. The polymer viscosity is reduced in-flight by heating and melting, and the particles "splat" against the substrate. This environmentally compliant coating application technique does not require solvents because the starting material is in the form of a powder. Other important advantages of thermal spraying over other coating processes (such as sol-gel, fluidized bed, or vapor deposition processes) are that the deposition of nanoreinforced composites is not limited by the size of the part to be coated, and that the coatings can be readily applied in the field. One of the most important applications for thermally sprayed coatings is their use as corrosion protection coatings for metallic substrates. Previous results 10 demonstrated that the high velocity oxy-fuel (HVOF) combustion spray technique is also a viable method for the processing of nanoreinforced polymers.

The processing, physical properties, and microstructures of thermally sprayed silica or carbon black/nylon11 nanocomposite coatings have already been evaluated as a function of processing conditions and discussed in a previous publication. 11 It was demonstrated that combusting gas mixtures with a low hydrogen content were needed for optimal jet temperature, resulting in better particle flow and improved filler distribution in the polymer matrix. Thermally sprayed nanocomposite coatings exhibited an increase in crystallinity relative to unfilled coatings because of the presence of the filler: up to a 15% and up to a 9% increase in coatings produced from 60 μm and 30 µm polymer particles, respectively, relative to nonreinforced thermally sprayed polymer. Coatings produced from smaller polymer particles also had higher densities and lower porosity.

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THEORY

Extensive development of theories explaining reinforcement mechanisms and for the prediction of the properties of reinforced composites abound. 12-34 Application of these to nanocomposite properties due to the addition of nanofillers is not yet as well established.

A very general approximation of the behavior of filled polymers is the rule of mixtures, which provides upper and lower boundaries to mechanical response. The upper bound eq. (1) reflects strong adhesion between the polymer and filler and a high filler aspect ratio, whereas the lower bound eq. (2) is more applicable to rigid spherical particles:

$$M_c = M_p \phi_p + M_f \phi_f \tag{1}$$

$$M_c = \frac{M_p M_f}{M_p \phi_f + M_f \phi_p} \tag{2}$$

where M_c is the modulus (shear, elastic, or bulk) of the composite, M_p and M_f are the moduli of the polymer matrix and filler, and ϕ_p and ϕ_f are the volume fractions of the polymer and filler, respectively. The moduli of most particulate filled polymers have been shown to lie somewhere between these two limits. ¹² The main reinforcing mechanism was assumed to be the effectiveness of load transfer from the particles to the polymer matrix.

Particle geometry, size, degree of dispersion and agglomeration, and interfacial interactions strongly affect the final properties of filled polymers. Simple theoretical models describing the reinforcement of a material, such as the one proposed by Einstein¹³ and later extended by Guth and Smallwood,¹⁴ assumed perfect interfacial adhesion. These are valid up to low filler volume fractions, because they ignore any mechanical interaction between neighboring particles. A relation that also accounts for the effect of adhesion efficiency between the two phases has been suggested by Sato and Furukawa. 15 The models proposed by Mooney, 16 Eilers and van Dyck, 17 and Bills et al. 18 took into consideration a number of effects from the filler distribution. Experimental deviations from the theoretical predictions of these models due to particle size were, however, reported. 19 Quemada 20 introduced a variable coefficient to account for inter-particle interactions and differences in particle geometry. Frankle and Acrivos²¹ introduced the concept of maximum

packing fraction to account for differences ticle geometry.

Kerner's equation²² was derived to call the modulus of a composite containing a spherical particles with some adhesion be the phases at up to moderate concentrat Nielsen's modification of Kerner's equation more versatile at low reinforcement concentions, taking into account both the maxim packing fraction of the filler, inter-particle actions and the relative modulus of the two stituents²³:

$$\frac{M_c'}{M_p} = \frac{1 + AB\phi_f}{1 - \psi B\phi_f}$$

where A accounts for the contribution of polyfiller interactions and factors such as the generalized Einstein coefficient k_E (tabulate ref. 12) from:

$$A=k_E-1$$

 ψ in eq. 3 is a reduced concentration term ded dent on the maximum packing fraction ϕ_m of filler in the polymer (also tabulated in ref. 12)

$$\psi = 1 + \left[(1 - \phi_m)/\phi_m^2 \right] \phi_f$$

The constant B takes into account the relationed moduli of the filler and the matrix; its value is for very large M_f/M_p ratios. It is defined as:

$$B = \frac{M_f/M_p - 1}{M_f/M_p + A}$$

The principle of viscoelastic correspondence plants the substitution of storage or complex multi into eq. (3). The Einstein coefficient k_E is dispersed spheres with perfect adhesion to a mutrix, estimated by Lewis and Nielsen, ²⁴ has value of 2.5. The value drops to 1.0 if therein perfect slippage at the interface. For large-scal agglomeration, or in the case of elongated particles, the Einstein coefficient and the value of are expected to be higher (>3) than for well dispersed spherical particles. ¹²

Filler size effects were not directly considered in traditional reinforcement models. As shown by Boluk and Schreiber, 25 however, experimentally measured moduli of nanofilled polymers closely file.

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those calculated according to the Nielsen model. The modulus of thermally sprayed coatings will be calculated using this model.

The modulus of polymers also increases rapidly with crystallinity, especially above T_g . Crystallites are expected to act as crosslinks by tying segments of many molecules together. Furthermore, crystalline regions have higher moduli than amorphous ones, and as a result can behave as rigid filler in an amorphous matrix. To a first approximation, the modulus above T_g is related to the degree of crystallinity, expressed as a weight fraction w_c , by eq. (7) proposed by Nielsen and Landel¹²:

$$\log_{10}G \cong 5.763 + 4.77w_c \tag{7}$$

The comparison of the predicted modulus based in the amount of crystallinity with the experimental data can therefore be used to elucidate the of the possible reinforcing mechanisms.

Mechanical damping is also affected by the presence of fillers and by filler/matrix interface conditions. Damping behavior is particularly interesting around the T_g because of the changes accurring there due to the filler. As shown by Nielsen, 23 in filled polymers damping behavior is related to the polymer volume fraction:

$$\tan \delta_c = \tan \delta_p (1 - \phi_f) \tag{8}$$

There ϕ_f is the volume fraction of the filler, and δ_c and tan δ_p are the loss moduli of the composite and polymer, respectively. This equation is filled for composites filled with nonagglomerated apid solid particles and without interactions.

Significant physicochemical interactions or mads between the polymer and the filler can condute to the formation of an interface with distinctive behavior, 26 induce crystallization or orientation the chains in the immediate vicinity of the filler face, 27 or result in the formation of a "bound" or mobile polymer layer. 28 According to Boluk and direiber, 25 eq. (8) can be rewritten, with the introvion of a correction parameter B, as originally gested by Iisaka and Shibayama²⁹:

$$\tan \delta_c = \tan \delta_p (1 - B\phi_f) \tag{9}$$

correction parameter B is related to the eftive thickness of the particle-matrix interface through:

$$B = (1 + \Delta R/R)^3 \tag{10}$$

where R is the mean radius of the dispersed particles, and ΔR is the thickness of the immobilized layer. This hypothetical boundary layer in the presence of a filler is considered, by several authors, to be another possible reinforcing mechanism beyond the traditional load transfer, especially in nanocomposites where the reinforcing effect of the filler is unusually large even at low filler contents. ^{25–29} Predictions calculated according to this model will be used for comparison with results of this work.

One strong indicator of the corrosion resistance of a coating is its resistance to water permeation. Predictions of permeation processes in heterogeneous systems are as complex as the mechanical responses; the overall transport depends on the amount, distribution and nature of the components, the magnitude of component interactions, and any interfacial phenomena. Permeation through a polymer is governed by two intrinsic factors: polymer chain segment mobility and defect structures, such as micro-cracks, pores, and variations in density.30 The presence of regions which are inaccessible to the diffusing species within a polymer effectively increase the diffusion path length and tortuosity and reduce its permeability. A further decrease in diffusion coefficient has been observed with increasing crystallinity by the same process. 31,32 In the case of plate or flakeshaped particles diffusion depends strongly on their orientation.33 The permeability of polymers filled with flaky or lamellar particles is usually lower than those filled with spherical fillers.34

The quantification of changes in the permeation resistance associated with the heterogeneities in a continuous polymer medium is not straightforward, but some simplified approaches have been developed. Permeability of polymers containing an impermeable phase such as fillers or impermeable polymer crystals in the absence of porosity can be estimated from the model proposed by Barrer³⁵:

$$P_i = P_0 (1 - \nu_i)^2 \tag{11}$$

where P_i is the permeability of polymer containing an impermeable phase, P_0 is the permeability of an amorphous or unfilled polymer, and v_i is the volume fraction of the impermeable phase. This model is used to predict permeability for nanocomposite coatings in this work and the results are compared with the experimental data.

Table I Nominal Vs. Actual Filler Contents in Composite Powders and Fill d Nylon 11 D-30 and D-60 Coatings

5 2.4 3.2 — 10 4.5 6.2 4.6 15 6.7 7.7	Carbon D-60 C (vol	Silica in D-30 Coatings (vol %)	Silanated Silica in D-60 Coatings (vol %)	Silica in D-60 Coatings (vol %)	Nominal Filler Content in Powder (vol %)
10 4.5 6.2 4.6			3.2	2.4	5
15 67	4	. 46		4.5	10
	7	6.9	7.7	6.7	15
20 6.9	10	6.9		6.9	20

EXPERIMENTAL

Materials

Nylon 11 ("French Natural ES," Elfatochem North America, Inc., King of Prussia, PA) was chosen as the matrix material because it has a high chemical resistance and wide processing window-large difference between the melting (\sim 183°C) and degradation temperatures (\sim 360-550°C). The T_g is typically 53°C, varying slightly as a function of reinforcement and crystallinity. Nylon 11 has been widely used as a coating because of an excellent combination of properties, such as low-temperature flexibility, low coefficients of friction, superior mechanical strength, and high chemical resistance. In this study, nylon 11 powders with mean particle sizes of 30 and 60 μ m, designated as D-30 and D-60, respectively, were used.

Silica powders, 7 and 12 nm ("R 812" and "A 200," from Degussa Corporation, Ridgefield Park, NJ), with hydrophobic and hydrophilic surface chemistries, respectively, 5 µm precipitated silica, 6 nm carbon black (Degussa Corporation) and A 1100 gamma-aminopropyltriethoxy silane modified silica, referred to as "silanated silica," were used for nylon 11 reinforcement to determine the effect of the different surface chemistries and powder sizes on the nanocomposite properties. Silanes were applied to decrease or remove any hydrophilic properties of the silica surface, to reduce agglomeration, and to incorporate new organofunctional groups with higher chemical affinity to specific polymers on the SiO₂ surface.³⁶ Silica was mixed for 3 h in a silane-toluene solution at 5 wt % of silica in toluene and then dried at a temperature 105°C to remove the solvent. Coatings with nominal 10 vol % of precipitated silica with a mean particle size of 5 µm were also produced to show the effect of particle size on the coating properties.

Co-spraying two different powders can in significant segregation in the jet, because differences in powder size and density. A compite powder, containing both materials in a simpowder, aids both in simultaneous powder ing into the HVOF spray jet and in distributing the filler in the coating. Nylon 11 powder was ball-milled together with the nanoparticular phase for 48 h in a Norton Ball Mill using zirch balls to create the composite powder.

The nominal content of filler in the power compositions sprayed ranged from 5 to 15 vol. Table I shows the actual filler content in nanocomposite coatings, determined by therm gravimetric analysis as described below. In the discussion of the results, except where noted specimens are labeled according to the nominary volume percent of filler in the starting material.

Coating Procedure

A Stellite Coatings' Jet Kote II® HVOF combilition spray gun with internal powder injection and a 0.076 m (3") long and 0.008 m (5/16") diameter nozzle, was used to spray the composite powder Typical processing parameters, detailed in a privious publication, 11 are listed in Table II.

The nanosized silica-reinforced nylon 11 composites were deposited onto $25.4 \times 75.2 \times 3$ min $(1'' \times 3'' \times 0.125'')$ aluminum (6061) substrates and also 60 mm diameter aluminum (6061) disc. Before spraying, the substrates were grit-blasted using $1600~\mu m$ SiC grit and cleaned in an ultrasonic ethanol bath. Substrates to be used to support coatings for subsequent thermal and structural analyses were polished instead of gritablasted to allow for easy removal of the coating and the substrated of the coating of the substrated to allow for easy removal of the coating of the substrated to allow for easy removal of the substrated to allow for easy removal of the substrated to the substrated to allow for easy removal of the substrated to the sub

The substrates were preheated to approximately 80°C by traversing the HVOF jet over the substrate surface before powder injection. Typical coating thicknesses ranged between 250 to 350 μ m (0.01–0.014 in.).

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Table II HVOF Processing Parameters Used for Spraying of Polymer Nan composite Coatings

Parameter Total gas flow rate Hydrogen/oxygen ratio Hydrogen/oxygen gas pressure Powder feed rate	SI Units	Standard ^a Unit	
Total gas flow rate	$14.9-17.0 \times 10^{-3} \text{ m}^3 \text{ s}^{-1}$	1600–1820 scfh	
Hydrogen/oxygen ratio	0.29-0.50	_	
Hydrogen/oxygen gas pressure	0.83 MPa	120 psi	
Powder feed rate	0.25 g s^{-1}	15 g min ⁻¹	
Powder carrier gas	Nitrogen		
Powder carrier gas flow rate	$0.5 \times 10^{-4} \text{ m}^3 \text{ s}^{-1}$	60 scfh	
Powder carrier gas pressure Sample velocity	0.97 MPa	140 psi	
Sample velocity	Stationary		
Gun horizontal surface speed	0.23 m s^{-1}	45 ft. min ⁻¹	
Step size per pass	$3.2 \times 10^{-3} \text{ m}$	0.126 in.	
Gun horizontal surface speed Step size per pass Spray distance	0.2 m	8 in.	

^a Traditional thermal spray industry standard units.

Coating Characterization

The "as-sprayed" coatings exhibited peak-to-valley roughnesses up to 30 μm and therefore before scratch and wear testing the coating surfaces were polished using SiC paper to a roughness of <6-μm.

Scratch tests were performed using a BYK Gardner SG-8101 balance beam scrape adhesion and mar tester according to ASTM D 5178-91³⁷ at applied loads of 2, 2.5 and 3 kg. Scratch profiles and scratch depths were measured using a Hommelwerke model Dektak II stylus tracing profilometer. A typical scratch profile is shown in Figure 1. Scratch results represent the mean of nine measurements performed on three different specimens.

Sliding pin-on-disk wear tests were conducted using an AMTI Model C tribometer, according to ASTM G 99-90, using a 10 mm diameter 52100 stainless steel ball as a counterbody. The coatings were tested at room temperature in ambient air 0r 15,000 cycles, with a 10N applied load, at a furface sliding speed of 0.68 m s⁻¹. Wear track goss sectional areas were measured using the Hommelwerke profilometer. The track profile was measured at four points (Figure 2) on two wear discs with the same type of coating. Results were in an ultiliamean of eight measurements.

The coefficient of friction μ was calculated from be wear data using:

$$\mu = \frac{F}{L} \tag{12}$$

ection. Type here F is the friction force (N) and L the normal n 250 to (N). The data represent mean coefficient of nction under steady-state wear conditions.

Dynamic mechanical analysis was performed using a Rheometrics Scientific ARES analyzer operated in torsion mode. The in-phase and out-ofphase responses were measured at a frequency of 1 Hz over a temperature range of 20 to 150°C on 5-mm-wide strips of coatings.

The permeability of thermally sprayed coatings was measured using the permeability cup method (ASTM D1653-9338) according to Test Method A, under Condition A (Figure 3) at 50% relative humidity and at 23°C over a 21-day period. Under steady state conditions, the increase in weight with time corresponded to the amount

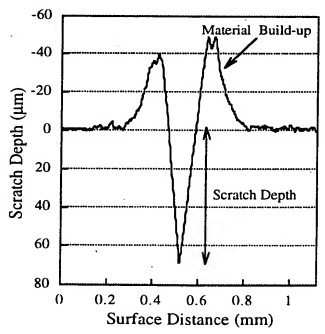


Figure 1 Typical scratch profile.

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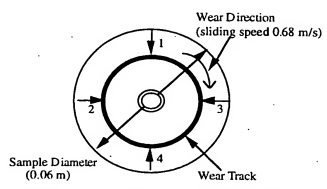


Figure 2 Locations of the wear track and profile measurements.

of water vapor diffusion. At least three testing cycles were performed for each coating composition.

Standard deviations are shown as error bars for the data obtained from averaging statistically significant numbers of measurements (at least five); otherwise a data range is reported.

RESULTS AND DISCUSSION

Scratch Resistance

Scratch resistances of 15% D-60 sprayed coatings at 2-kg applied load are shown in Figure 4. With an increase in reinforcement content the scratch depth of all composite coatings decreased. A maximum scratch resistance was achieved for coatings of either 15% hydrophobic silica or carbon black particles in a nylon 11 D-60 matrix at an applied load of 2 kg (Fig. 4). These increases represented 35 and 30% improvements in scratch resistance of nanocomposite coatings over pure nylon 11 D-60 coatings, respectively.

Scratch depth increased with increasing applied load for all samples (Fig. 4) but an increase

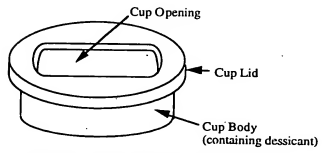


Figure 3 Permeability testing cup assembly.

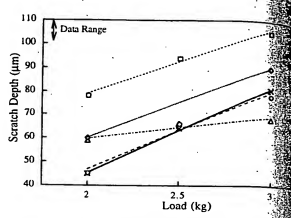


Figure 4 Scratch depth of thermally sprayed in 11 D-60 coatings containing: (\Box) 0, and nominal 18% of: (\bigcirc) hydrophobic silica, (\diamond) hydrophilic silica silanated silica, and (\times) carbon black fillers.

in reinforcement similarly decreased the scridepth. Fifteen percent silanated silica nanog posites exhibited the lowest scratch depths of samples at 3 kg load, a 35% decrease relative pure thermally sprayed nylon 11 D-60. Silana silica filled coatings also did not exhibit as gran increase in scratch depth with load as untreated silica coatings. A more even dispersion of the reinforcing particles in the silanated silical ples¹¹ was likely the cause of this behavior.

The scratch depths of D-60 coatings reinforce with micron-sized particles applied by the fluid ized bed process were measured to be 57, 60, at 63 μ m at 2, 2.5, and 3 kg loads, respectively. These were comparable to thermally sprayed inforced D-60 coatings.

The scratch depth of the D-30 thermal sprayed composite coatings decreased slightly with increasing reinforcement content relative pure D-30 coatings (Fig. 5). This decrease was no as pronounced, however, as in the case of the D-6 composite coatings (Fig. 6). The differences scratch depth were closely related to the coating microstructure and polymer crystallinity (Fig. 7) The crystallinity content dominated the scrate resistance at lower loads (2 and 2.5 kg). This result was consistent with known improvement in hardness and mechanical properties of poly mers such as elastic modulus and yield stres with increasing crystallinity. 39,40 Figures 6 and show that particle/matrix interfacial interaction had a significant effect on both the scratch behav ior and the degree of crystallinity in the D-60 nanocomposite coatings. Figures 6 and 7 also

Figure 5 [1] D-30 cc 4 of (O) 1 fillers.

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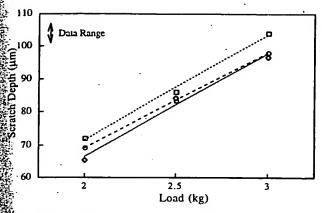
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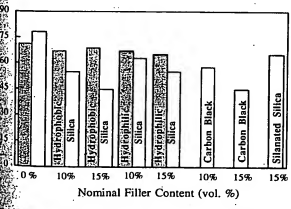
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show that 15% hydrophobic silica and 15% carbon black reinforced composites both exhibited identical scratch resistances, despite large differences in crystallinity.

The improvement in scratch resistance for the D-30 coatings as a function of crystallinity is shown in Figure 8. A comparison of Figures 7 and 8 indicated a greater increase in crystalline content in the D-60 coatings than in the D-30 composite coatings.

It was hypothesized that efficient load transfer was further affected by the degree of particle/inatrix interactions. These interactions depend on various factors, including the chemistry of the reinforcement surface, the efficiency of coupling



Mire 6 Scratch depth of thermally sprayed coateas a function of filler type and content at 2-kg load. Wand white bars represent D-30 and D-60 nylon 11 fices, respectively.

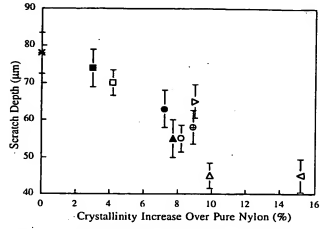


Figure 7 Scratch depth of thermally sprayed nylon 11 D-60 coatings as a function of filler and crystallinity content. The symbols represent: $(\Box, \bigcirc, \triangle)$ hydrophobic silica; $(\blacksquare, \bullet, \blacktriangle)$ hydrophilic silica; (\triangleright) silane treated silica; (\oplus, \triangleleft) carbon black. The star, square, circle, and triangle symbols represent 0, 5, 10, 15 vol % nominal filler contents, respectively.

between the particle and the matrix, and on the reinforcement dispersion. In Figure 9(a,b), typical scratch microstructures are shown. Figure 9(a) shows a scratch in an unreinforced D-60 coating where shearing of the polymer matrix along the whole length of the scratch is clearly visible. Numerous sheared, torn, and uneven surfaces appeared as white marks in the scratch valley. In a reinforced coating, as shown in Figure 9(b), shearing was greatly reduced. Damage to the matrix appeared to be localized to the polymer-rich areas

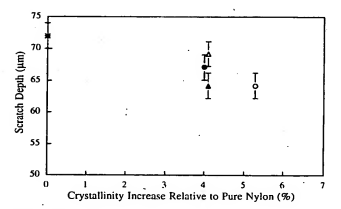


Figure 8 Scratch depth as a function of filler and crystallinity content for D-30 coatings. Legend: (*) 0, (\bigcirc) 10, (\triangle) 15 vol % of hydrophobic silica, and (\bullet) 10, (\triangle) 15 vol % of hydrophilic silica nominal contents.

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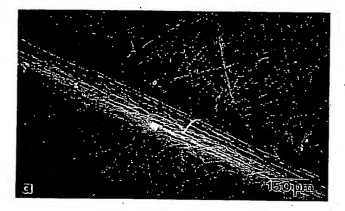
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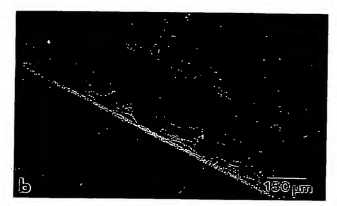


Figure 9 SEM micrographs of scratch surfaces of (a) an unfilled D-60 thermally sprayed nylon 11 coating, (b) a nylon 11 D-60 coating with nominal 10 vol % hydrophobic silica content.

inside "the cell" (as described previously¹¹), seen as repeating valleys along the scratch. Only a few sheared layers appeared to have been separated from the polymer matrix. The reinforced areas, consisting of embedded particles on the polymer particle surface, exhibited greater resistance to scratch penetration and material damage than nonreinforced areas. In the case of nonreinforced coatings, the scratch valley was more acute and deeper than that of reinforced coatings.

Results indicated that filler content and crystallinity had a dominant effect on scratch resistance, with an additional dependence on reinforcement distribution and chemistry.

Sliding Wear Resistance

Figure 10 shows the effect of reinforcement on the wear resistance of D-60 coatings. Wear decreased rapidly for all types of coatings with inclusion of

the reinforcement. The largest improvement and 67% wear track area reductions, occur 15% hydrophobic silica and 15% carbon coatings, respectively. Large differences in performance were found for different types of ticles and between samples with different crainity and filler contents, similar to the bendescribed for scratch resistance. The wear tance of a thermally sprayed nylon 11 contents are inforced with a nominal 10 vol % of 5-uniticle size precipitated silica is included for parison. As shown in Figure 10, this exhibit 13% wear track area reduction, but the reinforment was significantly less effective than an the nanoreinforcements at an equivalent loss.

The results in Figure 11 show again that ings with the highest filler and crystallinity tents, 15% hydrophobic silica and 15% carblack composites, exhibited the highest weak sistance. In the case of 15% hydrophobic silication 48% decrease in wear track area was measured the largest improvement in wear performant 67% decrease in wear track area, was measured for 15% carbon black filled coatings. The results for wear performance were consistent was scratch resistance results.

Figure 12(a,b) shows micrographs of typ wear tracks for pure and reinforced D-60 dings. The wear surfaces showed wear sheet mation and delamination in the sliding direction. The damage in the wear tracks of reinforced dings was significantly reduced [Fig. 12(b)], though the wear mechanism appeared to be a significant to the significan

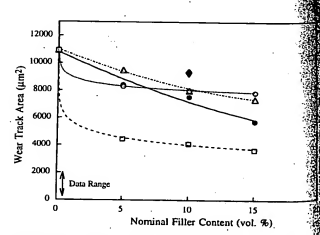


Figure 10 Wear track area of thermally sprayed nylon 11 D-60 coatings as a function of filler content. Symbols represent: (O) hydrophilic silica, () hydrophilic silica, (A) silanated silica, (\spadesuit) 5 μ m precipitated silica, and (\square) carbon black fillers.

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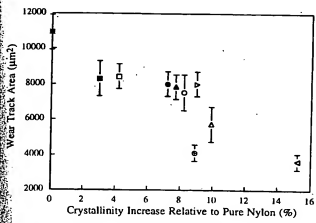
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Figure 11 Wear track area of D-60 nanocomposite coatings as a function of filler and crystallinity content. Symbols represent: (□, ○, △) hydrophobic silica, (■, ●, ♠) hydrophilic silica, (▷) silane treated silica, (⊕, ▷) carbon black. 0, 5, 10, and 15 vol % nominal filler contents are represented by star, square, circle, and friangle symbols, respectively.

same as for the pure polymer coatings. No wear craters were observed in the wear-damaged area, which would be characteristic of particle pull-out. The greater wear track area reduction in carbon black reinforced coatings may also have been due to the so-called "self-lubricating" nature of the arbon black particles.

The results of wear testing of D-30 coatings are hown in Figure 13. The wear track area of these hermally sprayed nanocomposite coatings deceased with the addition of reinforcement by up 25% relative to pure D-30 polymer coatings. The largest reduction in wear track area was discreted for 10% hydrophobic silica filled coatings. D-30 coatings exhibited lower wear resistance than equivalent D-60 coatings, similar to behavior observed for scratch resistance. It is was most probably due to the lower restallinity contents of the D-30 coatings.

Coefficients of friction (μ) of the sprayed coatgraph are summarized in Table III. The coefficients friction of D-60 reinforced coatings increased with the addition of reinforcement related to the pure nylon D-60, whereas the coeffient of friction of the D-30 nanocomposite coatgraph remained nominally the same as that of the trep-30 polymer coatings. An increasing μ with reasing reinforcement content was also reted for high density polyethylene filled with the gel. 42 Ramasubramanian et al. 42 reported linitial increase in μ , followed by a decrease for



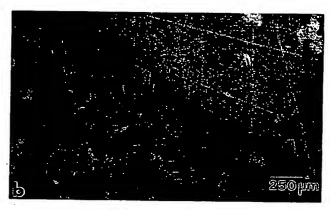


Figure 12 SEM micrographs of wear track surfaces of (a) an unfilled thermally sprayed nylon 11 D-60 coating, (b) a nylon 11 D-60 coating with nominal 10 vol % hydrophobic silica content.

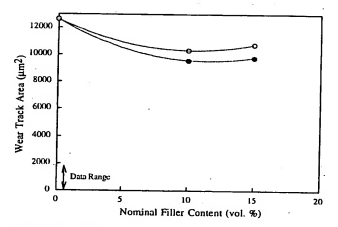


Figure 13 Wear track area of nylon 11 D-30 coatings as a function of filler content. Symbols represent: (O) hydrophilic silica and () hydrophobic silica fillers.

Table III Coefficients of Friction of Thermally Sprayed Nanocomposite Coatings

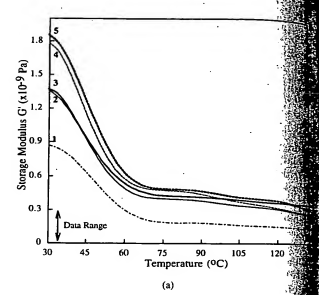
Silica Content (Nominal Vol %)	Silica Surface Chemistry	Nylon 11 Matrix	Mean Coefficient of Friction
0	_	D-60	0.24
0		D-30	0.29
5	Hydrophobic	D-60	0.24
5	Hydrophilic	D-60	0.27
5	Silanated	D-60	0.25
10	Hydrophobic	D-60	0.28
10	Hydrophilic	D-60	0.25
10 '	Silanated (5 µm)	D-60	n/a
10	Precipitated	D-60	0.33
10	Hydrophobic	D-30	0.29
10	Hydrophilic	D-30	0.28
15	Hydrophobic	D-60	0.30
15 -	Hydrophilic	D-60	0.29
15	Silanated	D-60	0.25
15	Hydrophobic	D-30	0.28
15	Hydrophilic	D-30	0.28

graphite and molybdenum disulfide filled polyethylene at higher filler contents. This critical content seemed to be dependent on the type of filler used and the testing conditions. Clerico⁴³ reported that the coefficient of friction decreased only when a polymer transfer layer was formed on the metal counterbody, thus changing the contact conditions. The addition of a larger size filler resulted in a larger increase in μ than was observed for nanoreinforcement, as is more typically expected from filled coatings. ^{42,43}

Dynamic Mechanical Properties

A series of representative storage modulus (G')curves as a function of temperature, recorded at 1 Hz for both D-30 and D-60 coatings containing 0-15% of silica are shown in Figure 14(a) and (b), respectively. The storage moduli both below and above the T_{g} increased with increasing amounts of silica reinforcement. The curves for the D-30 coatings [Fig. 14(a)] clearly show the region associated with the T_g . A clear transition was also observed for D-60 coatings [Fig. 14(b)]. In the case of D-60 reinforced coatings, with a 120% higher modulus than pure polymer coatings, the glassy plateau and smaller decrease in slope of the G'curve relative to D-30 coatings indicated a higher crystallinity content in the D-60 coatings (also confirmed by differential scanning calorimetry

and X-ray analysis¹¹). For both the D-30 and matrices, hydrophobic silica reinforced can exhibited larger increases in dynamic modulus than coatings with the same contahydrophilic silica.



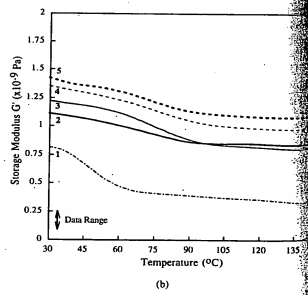


Figure 14 (a) Temperature dependence of the storage modulus G' of thermally sprayed D-30 nylon 12 coatings. Lines represent: (1) 0, (2) nominal 10, and (4) 15 vol % of hydrophilic silica; (3) nominal 10 and (5) 16 vol % of hydrophobic silica. (b) Temperature dependence of the storage modulus G' of thermally sprayed D-60 nylon 11 coatings. Lines represent: (1) 0, (3) 5, and (4) 15 vol % of hydrophilic silica; (2) 10 and (5) 15 vol % of hydrophobic silica nominal content.

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Table IV Dynamic Storage Modulus G' f Thermally Sprayed Nanocomposite Coatings

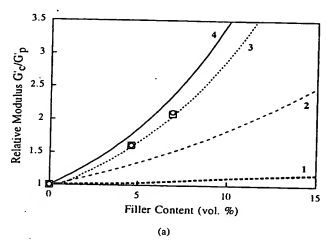
Silica Content (Nominal Vol %)	Silica Surface/Filler Type	Polymer Matrix	G' at 30°C (GPa)	<i>G</i> ′ at 70°C (GPa)
0	· _	D-60	0.65	0.36
0	_	D-30	0.86	0.35
<u>.</u> 5	Hydrophobic	D-60	1.15	. 0.86
5	Hydrophilic	D-60	1.58	1.07
5	Silanated	D-60	1.25	0.39
5	Carbon black	D-60	0.48	0.29
10	Hydrophobic	D-60	1.10	0.89
. 10	Hydrophobic	D-30	1.38	0.59
10	Hydrophilic	D-30	1.38	0.54
10	Silanated	D-60	1.29	0.61
ž 10	Carbon black	· D-60	0.50	0.32
15	Hydrophobic	D-60	1.39	1.06
15	Hydrophobic	D-30	1.83	0.61
15	Hydrophilic	D-60	1.31	0.99
15	Hydrophilic	D-30	1.79	0.56
15	Silanated	D-60	1.98	0.83
15	Carbon black	D-60	0.50	0.25

The dynamic storage moduli of thermally sprayed coatings at 30 and 70°C are summarized in Table IV. The largest increase, 205%, in the glass storage modulus was observed for the 15% filanated silica filled D-60 coatings, believed to be que to the improved particle distribution in the polymer matrix, in addition to the increase due to mystallinity. The glass storage modulus of D-60 poatings at 30°C increased by 120% and 100% elative to pure nylon 11 D-60 for hydrophobic and hydrophilic silica, respectively. The glass forage modulus of D-30 coatings at 30°C inreased by 182% and 175% for hydrophobic and ydrophilic silica, respectively, relative to pure ylon 11 D-30 coatings. The glass storage moduli D-30 coatings were typically higher, in the ange 0.86-1.83 GPa, relative to the D-60 coatngs, with moduli ranging between 0.65-1.39 Ta. This was believed to be the result of the improved filler distribution in the D-30 coatings ind more efficient load transfer between the polyer matrix and the reinforcing particles.

The opposite trend was observed for the storge modulus at 70°C. Both D-30 and D-60 reinticed coatings exhibited significantly higher oduli compared with coatings of the correspondpure polymer matrix. In the case of D-30, the mamic storage modulus was up to 70% higher, in the case of D-60 an increase of up to 195% the was believed to be due to higher crystallinity when the D-60 coatings.

Predictions of the nanocomposite moduli, calculated according to eqs. (1), (2), and (3), are shown in Figures 15(a) and (b) together with experimental values as a function of the reinforcement content. All the composite coatings exhibited a modulus higher than that calculated by the lower bound prediction (lines numbered 1). This may be due to particle/matrix and particle/particle interactions or variations in crystallinity. The solid lines 2, 3, and 4 were generated for A ranging between 2 and 4.5 with a maximum packing fraction $\phi_m = 0.37$, assuming a random close packing of agglomerated filler. The data for the D-30 silica reinforced coatings closely fit calculations for A = 3. For silica filled D-60 coatings the fit was closer for values of A between 3 and 4. There was a large deviation for the 5% filled D-60 coatings for both types of silica; the source of this deviation has not yet been determined. The values for A were in agreement with Nielsen's theory predicting higher values of both the Einstein coefficient and the constant A for agglomerated or elongated fillers; agglomerated particles were observed in the sprayed coatings, where the particles were concentrated along the splat boundaries forming a "layered" microstructure. It was believed that in the case of the D-30 coatings the lower values of A were due to an improved spatial distribution of the silica in the polymer matrix than in the D-60 coatings.

Equation (7) was used to estimate the effects of crystallinity increases on the moduli of sprayed



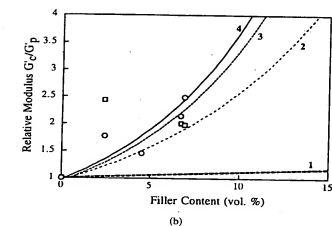


Figure 15 (a) Relative modulus G'
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otin G of nylon 11 D-30 coatings at 30°C. Symbols represent experimental values for: (O) hydrophobic silica, (\square) hydrophilic silica filler; lines are predictions according to: (1) the rule of mixtures, (2, 3, 4) Kerner's model for A=2, 3, 4, respectively. (b) Relative modulus G'
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nanocomposite coatings. The results are illustrated in Figure 16. The increases in moduli of nanocomposite coatings due to crystallinity increases was much higher than those predicted from the model. These estimates strongly indicated that other mechanisms, in addition to variations in crystallinity, were contributing to the reinforcement mechanisms in nanocomposite coatings.

 T_g is often reported as the temperature of the maximum loss tangent (tan δ). Values determined

in this manner are usually higher than sobtained from G'', because the temperature $(\tan \delta)_{\max}$ is much more sensitive to parameters such as crosslink density, filler content, of morphology than T_g itself. The damping per associated with the partial loosening of the permer structure, e.g., breaking of the intermolate bonds so that groups and small chain ments can move.

Over the 25 to 150°C temperature range stilled, one transition was observed for all compositions, corresponding to the T_g of the polymer matrix. T_g values, determined from the maimum in the loss modulus G'', are summarized. Table V. The values were similar to those tained from differential scanning calorimetric analysis. The addition of the reinforcement sulted in a 1–2°C increase in T_g for the Discoatings. Hydrophilic and hydrophobic silica fillid D-60 coatings exhibited a decrease in T_g with initial addition of reinforcement. At higher reforcement contents, however, an increase in T_g 3–4°C was observed. All silanated silica reforced coatings exhibited 1–2°C increases in T_g

Tan δ and its response to the presence of the silica reinforcement are shown in Figure 17(and (b) for D-30 and in Figure 18(a) and (b) for D-60 nylon 11 composites. The tan δ peaks at and 58°C corresponded to the maximum dampin of the D-60 and D-30 composites, respectively. The addition of nanosized fillers did not cause significant shifting of the damping peak; however broadening and lowering of the curve peaks with

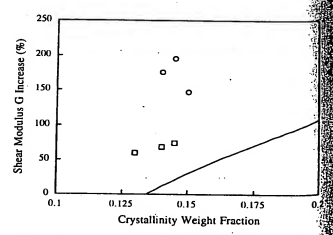


Figure 16 Shear modulus of nanoreinforced coatings as a function of crystallinity content. Experimental data are for: (O) D-60 and (\square) D-30 reinforced nylon 11 coatings. The solid line represents the shear modulus prediction according to eq. (7).

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Table V Glass Transition Temperatures (T_g) of Thermally Sprayed Nano omposite Coatings

Silica Content (Nominal Vol %)	Silica Surface Chemistry	Nylon 11 Matrix	T _g (°C)
0	•	D-60	50
0	_	D-30	48
5	Hydrophobic	D-60	46
5	Hydrophilic	D-60	47
5	Silanated	D-60	51
10	Hydrophobic	D-60	48
10	Silanated	D-60	52
10	Hydrophobic	D-30	50
10	Hydrophilic	D-30	50
15	Hydrophobic	D-60	54
15	Hydrophilic	D-60	53
15	Silanated	D-60	52
15	Hydrophobic	D-30	51
. 15	Hydrophilic	D-30	50

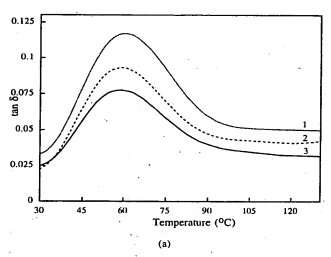
increased reinforcement contents was observed for both silicas in D-30 and D-60 coatings.

In Figure 19 (a) and (b), the damping behavior the D-30 and D-60 reinforced coatings, respecively, are compared with the predicted non-interctive damping behavior calculated according to fielsen's model [eqs. (9) and (10)]. According to his theory, deviations from non-interactive benivior suggest the existence of strong polymertemforcement interactions and potentially, the firmation of immobilized polymer layers in the ignity of the reinforcement particles. The values the layer thickness, ΔR , reported in the literatire vary, depending on the size and volume fracion of the filler particles and also on the polymer stem studied. Kendall and Sherliker²⁸ reported nm-thick "bound" polymer layer in polyethylfilled with nanosized silica and carbon black. dimilar results for the immobile layer thickness, the range of 0.5-2 nm, were reported by Brien et al.44 for carbon black in rubber. Iisaka hd Shibayama,²⁹ using eqs. (9) and (10), calcuted thicknesses up to 1.4 μ m for polystyrene/ ass beads (particle radii of $20-60 \mu m$) compos-Using the same model, Boluk and Schrecalculated the thickness of the immobile ger to be up to 20 nm in chlorinated polyethylfilled with nanosized rutile (TiO2). The presgof a layer with restricted mobility can result decrease in the loss tangent amplitude and a in the peak maximum to higher tempera-

The correction parameter, B, was evaluated the slope of the tan δ versus ϕ_f plots. The

thickness of the "immobilized polymer layer," ΔR , calculated from B values using eq. (10), are summarized in Table VI. The ΔR values, ranging from 3 to 23 nm, appeared to be within the range reported in the literature for various sizes of fillers. $^{25,27-29,44}$ ΔR values were lower, 3–7 nm for D-30, compared with 7–11 nm for D-60 coatings, probably again due to differences in the crystal-linity content of the two coatings.

Nanosized fillers contributed to crystallization in thermally sprayed nanocomposite coatings as reported previously¹¹ and therefore it is reasonable to suggest that some increase in crystallinity may be concentrated in the vicinity of the filler



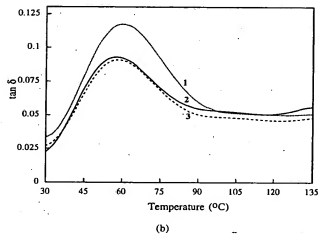


Figure 17 (a) Temperature dependence of tan δ for thermally sprayed nylon 11 D-30 coatings with: (1) 0, (2) 15, and (3) 10 vol % nominal hydrophobic silica content. (b) Temperature dependence of tan δ for thermally sprayed nylon 11 D-30 coatings with: (1) 0, (2) 15, and (3) 10 vol % nominal hydrophilic silica content.

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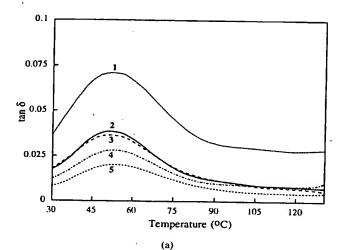
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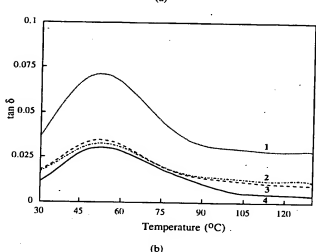


Figure 18 (a) Temperature dependence of tan δ for thermally sprayed nylon 11 D-60 coatings with: (1) 0, (2) 5, (3) 10, (4) 15, and (5) 20 vol % nominal hydrophobic silica content. (b) Temperature dependence of tan δ for thermally sprayed nylon 11 D-60 coatings with: (1) 0, (2) 20, (3) 15, and (4) 5 vol % nominal hydrophilic silica content.

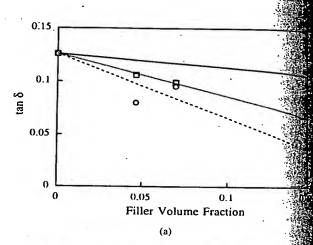
surface. The lamellar spacing of nylon 11 was reported to be between 7–11 nm, depending on the polymer processing and crystallization temperature, 45 consistent with the calculated ΔR values. The morphology of the "immobile" layer in thermally sprayed nanocomposite coatings, however, has not been confirmed at this stage.

Permeability

The results shown in Figure 20 indicated that the water vapor transmission rate (WVTR) through the thermally sprayed nanocomposite coatings

decreased with the addition of nanosized and carbon black reinforcements. Five phydrophobic and 5% hydrophilic silica reinforcements. D-60 coatings exhibited 16% and 18% decreased in WVTR, respectively, relative to pure D-60 mer coatings. With increasing reinforcement tents the WVTR decreased further, reaching maximum of 22.5% for 15% hydrophobic preinforced D-60 coatings. Only small different wVTR between different types of reinforced were observed.

More significantly, pure nylon 11 D-30 coat exhibited substantially lower values (by ~50



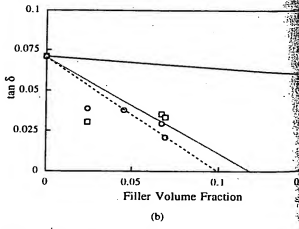


Figure 19 (a) Loss tangent as a function of filled content for nylon 11 D-30 coatings. Experimental data for (O) hydrophobic and (D) hydrophilic silica filler. This solid line is the prediction for rigid, non-interacting fillers [according to eq. (9)]. (b) Loss tangent as a function of filler content for nylon 11 D-60 coatings. Experimental data for (O) hydrophobic and (D) hydrophilic silica filler. The solid line is the prediction for rigid non-interacting fillers [according to eq. (9)].

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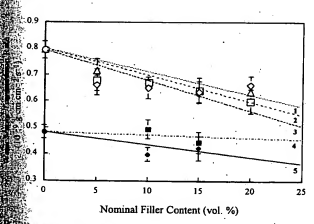
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Table VI Values of B Parameter and ΔR of Thermally Sprayed Nanocomposite Coatings Calculated According to Eq. (10)

Silica		<u>.</u> .		
Content (Nominal	Silica Surface	Polymer		ΔR
(Vol %)	Chemistry	Matrix	В	(nm)
5	Hydrophobic	D-60	18.6	11.5
5	Hydrophilic	D-60	23.5	22.4
5	Silanated	D-60	25.2	13.5
10	Hydrophobic	D-60	10.2	8.2
10	Hydrophobic	D-30	7.8	6.9
10	Hydrophilic	D-30	3.4	6.1
10	Silanated	D-60	13.3	9.6
15	Hydrophobic	D-60	8.7	7.4
15	Hydrophobic	D-30	3.5	3.6
15	Hydrophilic	D-60	7.4	11.4
黔 15	Hydrophilic	D-30	3.1	5.5
15	Silanated	D-60	10.7	8.4

WVTR than pure D-60 coatings. The incorporation of the silica filler in the D-30 coatings resulted in further significant decreases in WVTR. The lowest WVTRs were exhibited by D-30 coatings filled with 10 vol % of hydrophilic silica. Equivalent reductions in WVTR using hydrophobic silica in D-30 coatings required the addition of mominal 15 vol % content reinforcement.

Permeation of water vapor is believed to occur almost exclusively through the amorphous re-



The 20 Changes in WVTR of thermally sprayed accomposite coatings due to increasing amounts of increment. Legend: unfilled symbols, nylon 11 D-60 apposites; filled symbols, nylon 11 D-30 composites: 10° 0% reinforcement, $(\Box, \blacksquare, \text{lines } 1, 4)$ hydrophobic 10° 0, 10° 0, 1

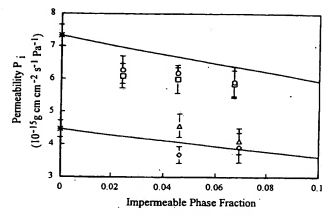


Figure 21 Permeability of nanocomposite coatings as a function of crystalline (impermeable) phase content. Experimental data are for: (○) hydrophobic and (□) hydrophilic silica fillers in D-60 nylon 11; (△) hydrophobic and (♦) hydrophilic silica filler in D-30 nylon 11. Solid lines represent the permeability predicted according to eq. (11).

gions of the polymer; a decrease in diffusion coefficient with increasing crystallinity is usually observed.³⁰ As expected, aqueous permeability decreased with both increasing crystallinity and reinforcement contents. The improvement was not linear with reinforcement content, showing the greatest changes after the initial addition of reinforcement, and leveling off at higher contents. This appeared to be a result of filler distribution; despite an increase in filler content, a significant volume of the filler remained concentrated in silica rich areas at the splat boundaries, 11 allowing increased diffusion through the polymer-rich areas. The initial addition of the reinforcing phase likely created the silica-rich barriers to vapor diffusion, therefore, such that additional filler had a less pronounced effect on the barrier properties.

As shown in Figure 21, the aqueous permeabilities of D-60 nanocomposite coatings were lower than predicted from theory according to eq. (11). Although the theory did not specifically separate the effects of crystallinity and filler, it was clear that the decrease was greater than that predicted for either effect alone, and that both effects likely were contributing simultaneously. The nylon 11 D-30 coatings, however, exhibited lower WVTR than D-60 coatings despite their lower crystallinity content.

As shown in Figure 22, the density of the coatings also appeared to have a significant effect on the coating permeability behavior. Coatings produced from the D-30 powders were always

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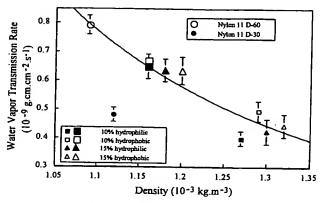


Figure 22 WVTR of thermally sprayed nanocomposite coatings as a function of density. Circle (\bigcirc) , square (\square) , and triangle (\triangle) symbols represent 0, 10, and 15 vol % nominal filler contents, respectively, in nylon 11 D-30 (smaller symbols) and nylon 11 D-60 (larger symbols). Open symbols (\square) and (\triangle) represent hydrophobic silica; closed symbols (\blacksquare) and (\triangle) represent hydrophilic silica.

slightly denser than those sprayed from the coarser D-60 powder. The nylon 11 D-30 coatings also exhibited significantly reduced WVTR compared with the D-60 with corresponding filler contents. Nylon 11 D-30 coatings also had slightly lower porosity contents, 0.5–1.5%, compared with 0.7–1.9% for D-60 coatings.¹¹

The slightly higher densities of the D-30 coatings were an expected result of the processing conditions: the smaller D-30 particles likely were more homogeneously heated than the larger D-60 particles. The more completely melted D-30 particles filled the underlying interstices more effectively upon splatting, thereby reducing the number and volume of pores created during deposition and increasing the cohesion of the coating. This led to a larger reduction in the WVTR for the D-30 coatings despite the higher crystallinity content of the D-60 coatings. Coating porosity and density therefore seemed to have a dominant influence on the aqueous permeability behavior of HVOF sprayed nanocomposite coatings with semicrystalline structures. Further detailed studies on the coating structure and crystal morphology are currently in progress to evaluate their effect on the coating permeability behavior.

CONCLUSIONS

Nanoreinforced polymer coatings deposited using the HVOF combustion spray process exhibited

significantly increased scratch and wea tance, and improved mechanical and properties over thermally sprayed pure policy coatings. Higher scratch resistances were sured in coatings produced from the powder larger polymer particles (D-60), likely due higher crystallinity of these coatings. The improvements in scratch resistance, 30 35%, were exhibited by 15% hydrophobic and 15% carbon black reinforced D-60 coats respectively. Similarly, the highest wear tances were measured for 15% hydrophobics and for all carbon black reinforced D-60 coal The increased wear resistance represented and 67% improvements relative to therm sprayed pure nylon D-60 coatings.

Increases in crystallinity and reinforcement content both resulted in increases in the storage modulus for silica reinforced coating greater increase in dynamic storage modifi than expected from the rule of mixtures was man sured for all silica-reinforced coatings. The imum increase in dynamic storage modulus belie the T_g relative to pure nylon 11 D-60 coatings. exhibited by 15% silanated silica D-60 coating (205%), followed by 15% hydrophobic silica forced D-30 coatings (182%). The largest increase in modulus in D-60 coatings was measured 15% hydrophobic silica coatings (120%). Polyme reinforcement particle interactions appeared dominate this improvement in moduli below Above T_{R} , D-60 nanocomposite coatings exhibited higher moduli than D-30 coatings due to higher crystallinity content of the D-60 coating

The WVTR of all reinforced coatings we lower than those of pure polymer coatings. The 15% hydrophobic silica reinforced D-60 coating exhibited a 22.5% decrease in WVTR, reaching value of 0.63×10^{-9} g cm cm⁻² s⁻¹. The WVTR D-30 coatings was even lower, due to the decreasin porosity. The lowest WVTR of 0.39×10^{-9} g cm⁻² s⁻¹ was exhibited by 10% hydrophilic silication of D-30 coatings, a 50% improvement wVTR relative to unfilled D-60 coatings.

Crystallinity and reinforcement content seement to have a dominant effect on the mechanical properties of thermally sprayed coatings, whereas coating density dominated the permeation behavior of nanoreinforced coatings.

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